Antidiabetic and Antioxidant Effect Combination Vasconcellea pubescens A.D.C. and Momordica charantia L. Extract in Alloxan-Induced Diabetic Rats

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ABSTRACT

Introduction: Mountain papaya (Vasconcellea pubescens A.D.C.) and bitter melon (Momordica charantia L.) fruit extract have total flavonoids and some metabolite from its contains. The use of a single dose in the treatment of diabetes is often considered to lack a strong effect. Giving a combination of the compound is a solution to provide a synergistic effect in treatment. Material and Methods: Rat were divided into eight groups (normal, negative control, glibenclamide as a positive control, and five dose extract group). The single dose mountain papaya extract (MPE) and bitter melon extract (BME) were given at 174 mg/kg b.w of MPE, and 380 mg/kg b.w of BME in oral administration. Combination of MPE:BME (25:75, 50:50, 75:25)% or (87:190; 44:285 and 130:95) mg/kg b.w in oral administration. Induction of diabetes used alloxan dose of 150 mg/kg b.w intraperitoneally. The treatment was performed for 21 days with a frequency of once a day. Blood sugar level was measured at pretest, 7th, 14th, and 21st days. At the end of the test, measurements of malondialdehyde (MDA), glutathione (GSH), and pancreatic morphology were measured. Data were analyzed by ANOVA statistic. Results: The single and combination of MPE-BME showed an decrease in blood glucose levels significantly compared with the negative control (p ≤ 0.05). The same results from measurements of levels of malondialdehyde (MDA), glutathione (GSH), and pancreatic morphology. Conclusions: Based on the research, a combination of the MPE-BME had antidiabetic and antioxidant activity, but the activity was not significantly different from both single-dose extract (p> 0.05).

Key words: Antidiabetic, Antioxidant, Flavonoids, Momordica, Synergistic, Vasconcellea.

INTRODUCTION

Diabetes mellitus (DM) is a metabolic disorder that occurs because the pancreas cannot produce insulin or the body is unable to use insulin produced effectively.¹-³ Indonesia estimates the 6th rank after China, India, the United States, Brazil and Mexico with 10.3 million sufferers. The use of oral antidiabetic drugs that are long enough for a patient’s lifetime is done only for them. Many studies have been conducted related to the use of oral antidiabetic drugs.⁴-⁶ Therefore, many people choose to use traditional medicine that is accepted to be safer, affordable, and effective.⁶-⁷

Mountain papaya (Vasconcellea pubescens A.D.C.) and bitter melon fruit (Momordica charantia L.) are several types of plants that are commonly found in Indonesia. Mountain papaya contains flavonoids, alkaloids, polyphenols, cysteine proteases, and papain which are antidiabetic agents.⁸-⁹ Total flavonoids contained in mountain papaya fruit have been permitted as antidiabetic by preventing β cell damage, insulin, and insulin signaling and inhibiting α-glucosidase.⁹,¹⁰ Bitter melon fruit contains flavonoids, alkaloids, saponins, polysaccharides, proteins, triterpenoids, quinine, and amino acids.¹¹ Fruit extracts can increase insulin secretion and protect the pancreas, inhibit α-glucosidase and inhibit α-amylase.¹²-¹⁴ Innovations in oral antidiabetic agents from natural ingredients have been developed. The use of a single dose in the treatment of diabetes is often considered to lack a strong effect. Giving a combination of ingredients is a solution to provide a synergistic effect in a treatment. In this study, we tested the antidiabetic activity of a combination of ethanol extract of mountain papaya and bitter melon fruit.

MATERIALS AND METHODS

Materials

Wistar rats were obtained from Faculty of Medicine, Universitas Sebelas Maret, Surakarta, Indonesia. Standard quercetin (Sigma Aldrich®), alloxan monohydrate (Aldrich®), reagent of lipid peroxidation and glutation colorimetric, aluminum chloride colorimetric reagent, glucose oxidase-phenom aminoantipyrine (GOD-PAP) from Biosystem, hematoxylin and eosin.

Extraction and analysis of total flavonoids

Mountain papaya and bitter melon extracted separately using the maceration method with ethanol 70% solvent. The test material obtained was washed and separated from the seeds, cut into small pieces and dried with a 40°C oven. Dry simplisia was macerated for 3x24 hours. The filtrate was evaporated with a rotary evaporator at 50°C until a thick extract was obtained.¹⁵ The method used is the aluminum...
cholorolimetric method using a UV-Vis spectrophotometer at λ = 439 nm. The standard used is Quercetin concentration series 6.125, 12.5, 25, 50, and 100 ppm.\textsuperscript{16}

**In vivo experiment**

The animal handling was approved from the health research ethics committee of Faculty of Medicine of Universitas Sebelas Maret No. 1046/I/IV/HEREC/2019. The rats were acclimatized for 7 days, then measured blood sugar levels pretest. Animal diabetes was done by inducing with alloxan monohydrate dose 150 mg / kg body weight (b.w) intraperitoneal (i.p).\textsuperscript{17} After 72 hours rats were treated with intensity one time a day. Animal experiments were divided into 8 groups: normal (I), positive control (II), negative control (III), and 5 test groups. Glibenclamide 0.45 mg / kg b.w was used as positive control. Test groups consist of combination MPE-BME extract 50%:50% (IV), 25%:75% (V), 75%:25% (VI), MPE 100% (VII), and BME 100% (VIII).

The dose of MPE 100% was 173 mg / kg b.w and the dose of BME was 100% 380 mg / kg b.w. The extract dose was based on the total flavonoid value of each extract.\textsuperscript{19} On the 21\textsuperscript{th} day euthanasia was performed with cervical dislocation, then pancreatic and hepatic organ harvesting was carried out.

**Blood glucose level**

Blood glucose measurements were made before alloxan induced (pretest), the day-7,\textsuperscript{10,14} and 21\textsuperscript{th}. Blood glucose measurement using GOD-PAP method with the Biochemisty Analizer by Biosystem instrument λ=500 nm.

**Rat liver homogenate**

Taken 500 mg of liver added in 5 mL of a solution of 0.15 M tris-HCl (pH 7.4) then shaken to homogenize it to obtain a homogenate of 10% b/v.\textsuperscript{19}

**Malondialdehyde (MDA) measurement**

Plasma malondialdehyde measurement was done using the thio-barbituric acid reactive substances (TBARS) method with a UV-Vis spectrophotometer. 0.2 mL liver homogenate, plus 0.2 mL sodium decylate sulfate (SDS) 8.1%, 1.5 mL of acetic acid 20%, 1.5 mL of TBA 0.8%, and aquadest and heated at 95°C for 60 minutes, then added 4 mL of 10% TCA, centrifuged at 3000 rpm for 10 minutes. After that the absorption is read at a wavelength of 532 nm.\textsuperscript{20,21}

**Glutathione (GSH) measurement**

Glutation level measurement refers to the method developed by Ellman et al., (1984), liver homogenate (0.75 mL) was added with 0.75 mL of 10% TCA, centrifuged at 200 rpm for 10 minutes, added with 1.8 mL ellman reagent (5, 5’-dithio bis-2-nitrobenzoic acid), absorbance was measured by UV-Vis spectrophotometer λ = 412 nm.\textsuperscript{22}

**Histological study**

Pancreatic histology testing aims to look at the general morphology of the pancreas. Fixation of pancreatic organs using 10% NBF solution. Tissue preparation of pancreas was colored with Hematoxylin-Eosin (HE). Hematoxylin would give a blue color on the nucleus and eosin gave a red color on cytoplasm and extracellular matrix.\textsuperscript{23} Histological examination carried out qualitatively to see the general morphology of pancreatic cells by counting the number of normal pancreatic cells and pancreatic cells undergoing necrosis in the form of nuclear shrinkage (pyknosis), nucleus rupture (cariorrexis), and disappearance of nucleus (karyolysis) on the Langerhans Island.\textsuperscript{9}

**Statistical analysis**

The analysis of blood sugar, GSH, and MDA levels was performed using statistics, while analysis of pancreatic histological data was carried out descriptively by looking at the general morphometry of pancreatic cells. Statistical analysis was preceded by a normality test with Shapiro Wilk and a homogeneity test with the Levene’s Test. Parametric tests were carried out with One Way ANOVA, post hoc LSD.

**RESULTS**

**Total flavonoid analysis**

The total flavonoid (quercetin equivalent) of MPE was 121.334 ± 3.404 mg / 100 g ethanol extract, while the total flavonoid of BME was 55.795 ± 1.601 mg / 100 g ethanol extract.

**Blood glucose analysis**

On the day 21\textsuperscript{th}, after treatment there was significant normalization and homogenization of fasting blood glucose every weeks, observed in diabetic experimental animals treated with combination MPE and BME. The standard as positive controls is glibenclamide 0.45 mg/kg b.w. Glibenclamid administer gave decrease fasting blood glucose that had p>0.05 from negative controls. MPE and BME have antidiabetic activities because they have significantly different sugar levels with negative controls. The antidiabetic activity of MPE and BME is almost the same (p>0.05) with glibenclamide. On the day 7\textsuperscript{th} the blood sugar levels of mice increased by 126.661-201.197 mg/dL. Blood sugar levels of rats after alloxan induction ranged from 200-260 mg/dL can be said to experience hyperglycemia.\textsuperscript{24} Blood sugar levels on the day 7\textsuperscript{th} of all extract treatment groups differed significantly from the negative group, but groups IV and VI did not differ significantly from positive controls. On the 14\textsuperscript{th} and 21\textsuperscript{th} days the blood sugar levels of the treatment group extract (IV-VIII) both combination and single were significantly different from negative control, but not significantly different from positive control (Figure 1).

**Histological analysis**

The treatment extract group that had the lowest total cell damage was group IV namely the test animals that were given CFE and BFE (50:50). This is thought to occur because the combination of a 50:50 MPE and BME ratio may has a synergistic effect in improving pancreatic morphology. The presence of flavonoid content in MPE and BME has antioxidant properties that are able to ward off free radicals causing pancreatic cell damage and inhibit pancreatic cell damage so that the extract treatment group has higher normal cell counts compared to negative controls.

**Glutathione (GSH)**

The results of normality and homogeneity tests showed that all groups in this study had normal and homogeneous data distribution with p values> 0.05 (Table 1). ANOVA results showed a significant difference

![Figure 1: Effect of combination ethanol extract (MPE-BME) on blood glucose of rats.](image-url)
between the negative group with the CFE and BFE groups ($p < 0.01$). The lowest GSH levels are the negative control group induced by alloxan, because alloxan can cause the failure of endogenous antioxidant defense mechanisms in counteracting free radicals. The lower the level of GSH, the higher the damage to the liver. GSH in the positive control group $31.8778 \pm 1.8558 \mu M/g$ of the liver showed that glibenclamide can suppress the hepatotoxic activity of alloxan. Groups IV, VII, and VIII had GSH values that were not significantly different from positive control ($p > 0.01$). The GSH value of the treatment group combination extract was not significantly different from the single extract treatment group (Figure 2).

**Malondialdehyde (MDA)**

The highest MDA level is known in the negative control group induced by alloxan. While the drug control group (glibenclamide) has the lowest MDA levels (Table 1). High levels of MDA indicate cellular damage due to free radicals. All extract groups (IV-VIII) were significantly different ($p < 0.01$) from negative control, but only groups IV and VIII were not significantly different from positive controls ($p > 0.01$). This shows that the combination of MPE-BME (50:50) and BME (100) has almost the same activity as glibenclamide because MPE and BME contain flavonoids which act as strong antioxidants compared to vitamin C and vitamin E. The MDA value of the MPE group (VII) was not significantly different ($p > 0.01$) from the BME group (VIII) and the MDA value of the combination group (IV-VI) was not significantly different ($p > 0.01$) from the MPE and BME groups (VII and VIII).

**DISCUSSION**

The antidiabetic activity of BME is higher than that of MPE because its fruit containing p-insulin polypeptide which works like insulin and charantin which can increase glucose in liver cells and muscles, has a component that works similar to sulfonylurea. Histology results of normal cell pancreas are shown with a rounded cell shape with a clear nucleus, uniform cell shape, and do not experience edema. The pancreatic tissue of the negative control group has more necrosis cells than the treatment of both single and combination MPE and BME extracts, which indicates more severe damage due to alloxan induction. Cells that experience pyknosis are essentially wrinkled and basophilic, dark in irregular boundaries. The cell nucleus that undergoes cariöerexis will fragment or disintegrate by leaving fragments of chromatin substances scattered within the cell. In the nucleic cell undergoing karyolysis, chromatin basophilic turn pale, the cell nucleus loses the ability to be stained and simply disappear. This condition is severe damage and irreversible even with cell regeneration activity.

Measurement of GSH levels is done by referring to the Ellman method. In principle, ditio bisnitro benzoate (DTNB) will react with GSH to produce a yellow tionitro benzoate compound. Malondialdehyde is a metabolite that results from lipid peroxidation by free radicals. The principle of the TBARS method is that MDA will form a complex with pink TBA in an acidic atmosphere.

**Table 1: Antioxidant activity of MPE-BME with the GSH and MDA parameters.**

<table>
<thead>
<tr>
<th>Groups</th>
<th>Mean GSH level ($\mu M/g$ Hepar ± SEM)</th>
<th>Mean MDA level ($\mu M/g$ Hepar ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I Normal</td>
<td>31.6036 ± 2.6248$^a$</td>
<td>0.0359 ± 0.0054$^{a,\ast}$</td>
</tr>
<tr>
<td>II Glibenclamide</td>
<td>31.8778 ± 2.9558$^{a,\ast}$</td>
<td>0.0260 ± 0.0057$^{a,\ast}$</td>
</tr>
<tr>
<td>III Negative Control</td>
<td>13.8151 ± 1.8581$^{a,\ast}$</td>
<td>0.1064 ± 0.0094$^{a,\ast}$</td>
</tr>
<tr>
<td>IV MPE-BME(50:50)</td>
<td>30.1434 ± 1.7470$^{a,\ast}$</td>
<td>0.0503 ± 0.0032$^{a,\ast}$</td>
</tr>
<tr>
<td>V MPE-BME (25:75)</td>
<td>23.8242 ± 1.6836$^{a,\ast}$</td>
<td>0.0697 ± 0.0167$^{a,\ast}$</td>
</tr>
<tr>
<td>VI MPE-BME (75:25)</td>
<td>22.5729 ± 1.5534$^{a,\ast}$</td>
<td>0.0591 ± 0.0075$^{a,\ast}$</td>
</tr>
<tr>
<td>VII MPE (100)</td>
<td>26.2291 ± 2.0982$^{a,\ast}$</td>
<td>0.0759 ± 0.0099$^{a,\ast}$</td>
</tr>
<tr>
<td>VIII BME (100)</td>
<td>26.2304 ± 1.1320$^{a,\ast}$</td>
<td>0.0467 ± 0.0087$^{a,\ast}$</td>
</tr>
</tbody>
</table>

(a) Significantly different from negative control $p < 0.05$ (b) significantly different from negative control $p < 0.01$ (c) significantly different from positive control $p < 0.01$ (*) significantly different from VII group ($p < 0.01$) (#) significantly different from VIII group ($p < 0.01$).
CONCLUSION
The single of MPE at dose 173mg/kg b.w, BME at dose 380 mg/kg b.w, and combination of both (50:50, 25:75, and 75:25) have an antioxidant and antidiabetic effect. The combination of MPE and BME has the same effective antidiabetic activity as glibenclamide. Antidiabetic and antioxidant activity of the combination extract did not differ significantly from the single extract.

CONFLICTS OF INTEREST
The authors declared no conflicts of interest

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REFERENCES
Sasongko, et al.: Antidiabetic and Antioxidant Effect Combination Vasconcellea pubescens A.DC. and Momordica charantia L. Extract in Alloxan-Induced Diabetic Rats

GRAPHICAL ABSTRACT

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